

PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION
International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

<p>(51) International Patent Classification 7 : D21C 9/00, D21H 11/20, D06M 13/192, 15/263</p>		A1	<p>(11) International Publication Number: WO 00/39389</p> <p>(43) International Publication Date: 6 July 2000 (06.07.00)</p>
<p>(21) International Application Number: PCT/US99/29884</p> <p>(22) International Filing Date: 16 December 1999 (16.12.99)</p> <p>(30) Priority Data: 09/222,372 29 December 1998 (29.12.98) US</p> <p>(71) Applicant: WEYERHAEUSER COMPANY [US/US]; P.O. Box 2999, Tacoma, WA 98477-2999 (US).</p> <p>(72) Inventor: JEWELL, Richard, A.; 5215 146th Avenue SE, Bellevue, WA 98006 (US).</p> <p>(74) Agents: CRAWFORD, John, M. et al.; Weyerhaeuser Company, Patent Dept. – CH2J29, P.O. Box 2999, Tacoma, WA 98477-2999 (US).</p>		<p>(81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).</p> <p>Published <i>With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.</i></p>	

(54) Title: **CARBOXYLATED CELLULOSIC FIBERS**

(57) Abstract

Carboxylated cellulosic fibers, comprising cellulosic fibers covalently coupled to a carboxylating agent through an ester bond, wherein the carboxylating agent provides a carboxyl group to the fibers, and wherein the carboxylating agent is a polycarboxylic acid having one carboxyl group separated from a second carboxyl group by either two or three atoms, wherein the carboxylated fibers have a water retention value greater than or equal to the water retention value of the fibers from which the carboxylated fibers are formed. Methods for producing the fibers and for producing fibrous products that incorporate the fibers are also disclosed.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav Republic of Macedonia	TM	Turkmenistan
BF	Burkina Faso	GR	Greece			TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's Republic of Korea	NZ	New Zealand		
CM	Cameroon	KR	Republic of Korea	PL	Poland		
CN	China	KZ	Republic of Korea	PT	Portugal		
CU	Cuba	LC	Kazakhstan	RO	Romania		
CZ	Czech Republic	LI	Saint Lucia	RU	Russian Federation		
DE	Germany	LK	Liechtenstein	SD	Sudan		
DK	Denmark	LR	Sri Lanka	SE	Sweden		
EE	Estonia		Liberia	SG	Singapore		

CARBOXYLATED CELLULOSIC FIBERS

Field of the Invention

The present invention is generally directed to cellulosic fibers and, more
5 particularly, to carboxylated cellulosic fibers and methods for their formation and use.

Background of the Invention

The tensile or sheet strength of fibrous products derived from cellulose fibers is
due in large part to attractive fiber-to-fiber interactions. These interfiber interactions
10 include hydrogen bonding interactions between fibers having hydrogen bonding sites.
For cellulose, hydrogen bonding sites primarily include the hydroxy groups of the
individual cellulose chains.

The present invention relates to increasing the strength of cellulosic fiber sheets
by incorporating carboxyl groups into cellulosic fibers from which the sheets are made.
15 In accordance with the present invention, carboxyl groups are incorporated into
cellulosic fibers through reaction with a carboxylating agent that is a polycarboxylic
acid.

Treating cellulosic fibers with polycarboxylic acids is known in the art. For
example, polycarboxylic acids have been used as crosslinking agents for cellulose.
20 Cellulose has been modified by reaction with dicarboxylic acids and their derivatives to
form simple diester crosslinks. Phthalic, maleic, and succinic anhydrides have been
used to form diester crosslinks in cellulose. Cotton has been treated with dicarboxylic
acid chlorides having varying chain lengths (e.g., from succinyl to sebacyl) to provide
ester crosslinks. Dicarboxylic acids have also been reacted with cellulose to provide
25 crosslinked cellulose containing diester crosslinks of various lengths (e.g., C₃-C₂₂).
However, oxalic acid has been shown to be unreactive to cellulose crosslinking, and
succinic and glutaric acids have been shown to have only slight reactivity. For a
review of ester crosslinked cellulosic fibers, see Tersoro and Willard, CELLULOSE AND
CELLULOSE DERIVATIVES, Bikales and Segal, eds., Part V, Wiley-InterScience, New
30 York, 1971, pp. 835-875.

Polycarboxylic acid crosslinked fibers and their preparation and use are also described in U.S. Patents Nos. 5,137,537; 5,183,707; and 5,190,563, issued to Herron et al. The Herron patents generally describe the preparation and use of individualized, polycarboxylic acid crosslinked cellulosic fibers having advantageous reduced water retention value properties. These fibers have a C₂-C₉ polycarboxylic acid crosslinking agent reacted with the fibers in the form of an intrafiber crosslink bond. The cellulosic fibers treated with the polycarboxylic acid crosslinking agents are cured at elevated temperature (e.g., about 190°C) to exhaustively couple the polycarboxylic acid to the cellulosic fibers through ester crosslinks. The C₂-C₉ polycarboxylic acid crosslinking agents include citric acid, 1,2,3-propanetricarboxylic acid, 1,2,3,4-butanetetracarboxylic acid, and oxydisuccinic acid, among others.

Polymeric polycarboxylic acids have also been used to crosslink cellulosic fibers. The use of polyacrylic acid crosslinking agents, including copolymers of acrylic acid and maleic acid, is described in U.S. Patent No. 5,549,791, issued to Herron et al. These polycarboxylic acid crosslinking agents were found to be particularly suitable for forming ester crosslink bonds with cellulosic fibers. Unlike some conventional crosslinking agents (e.g., C₂-C₉ polycarboxylic acids such as citric acid) that are temperature sensitive, polyacrylic acid is stable at high temperature and, therefore, can be subjected to elevated cure temperatures to effectively and efficiently provide highly crosslinked fibers. The Herron patent describes curing polyacrylic acid treated cellulosic fibers at about 190°C for about 30 minutes to form interfiber ester crosslinked bonds.

The mechanism of crosslinking paper with polycarboxylic acids has been described. See, Zhou et al., *Journal of Applied Polymer Science*, Vol. 58, 1523-1534 (1995). Brief thermocuring of paper treated with aqueous solutions of polycarboxylic acids provided paper having excellent wet strength through crosslinking. The effectiveness of a polycarboxylic acid to impart wet strength to paper was found to increase with increasing polycarboxylic acid functionality (i.e., number of carboxyl groups). Butanetetracarboxylic acid (BTCA) was found to be more effective than tricarballylic acid (TCA), which in turn was found to be significantly more effective

than succinic acid (a dicarboxylic acid). The excellent wet strengthening properties of polycarboxylic acids such as BTCA and TCA were determined to reflect the acids' ability to form multiple, reactive anhydrides during the curing reaction either directly, in the form of a dianhydride for BTCA, or in a successive, stepwise mode for BTCA 5 and TCA. For succinic acid, such a consecutive reaction is more difficult and reaction with succinic acid leads to a substituted cellulose having a considerable proportion of single carboxylic acid groups attached to cellulose through an ester link. Because the residual single carboxyl group reacts with cellulosic hydroxyl groups at a slower rate, succinic acid has been shown to be a poor crosslinking and wet strength agent for 10 paper. See Zhou et al.

The mechanism of polycarboxylic acid crosslinking of papers has been shown to occur in four stages: (1) formation of 5- or 6-membered anhydride ring from polycarboxylic acid; (2) reaction of the anhydride with a cellulose hydroxyl group to form an ester and link the polycarboxylic acid to cellulose; (3) formation of additional 5- 15 or 6-membered ring anhydride from polycarboxylic acids' pendant carboxyl groups; and (4) reaction of the anhydride with other cellulose hydroxyl groups to form ester crosslinks.

Reaction of paper with succinic acid at 150°C results in the formation of ester bonds or links, the number of which increases with curing time. A small amount of 20 crosslinking is observed, and the amount of crosslinking increases significantly with curing time and higher curing temperatures.

While polycarboxylic acid reaction with cellulose leads to substitution and crosslinking, only interfiber ester covalent bonds can support paper structure when wet. Because the ester links are water stable, the crosslinks prevent swelling of fibers 25 and thus may help hold the paper's fibers together. Although the introduction of carboxy groups into paper through esterification may affect some aspects of the paper's characteristics, the paper's primary wet strength results from the formation of interfiber ester covalent bonds. Both crosslinking and formation of interfiber ester covalent bonds are essentially the same chemical reaction. It can be seen that the 30 critical factors are whether the fibers are in contact with one another during curing and

the ability of the polycarboxylic acid to undergo more than one esterification reaction with cellulose hydroxyl groups.

Although the number of carboxyl groups incorporated into a paper treated with succinic acid can be high, the resulting paper has little wet strength. Because these 5 pendant carboxyl groups are largely incapable of further reaction with cellulose's hydroxyl groups to provide interfiber bonds or crosslinked fibers under normal curing conditions, most of these pendant carboxyl groups remain free. The mere presence of carboxylic acid moieties in a paper's cellulosic fibers does not impart wet strength to the paper.

10 However, cellulosic fibers modified to include carboxyl groups have been shown to impart strength to sheets in which the fibers are incorporated. More specifically, fibrous sheets incorporating carboxymethylated cellulose and carboxyethylated cellulose have been found to be relatively easily fibrilated or repulped and formed into sheets having superior strength properties. See U.S. Patent 15 No. 5,667,637, issued to Jewell et al., and references cited therein.

The wet strength of fibrous sheets made from carboxymethylated and carboxyethylated cellulose can be further increased by blending the carboxylated fibers with a wet strength resin, particularly a cationic additive. See, for example, U.S. Patent No. 5,667,637, and references cited therein. Generally, the addition of carboxyl 20 groups to cellulose is believed to enhance the efficiency of the wet strength resin by imparting wet strength to fibrous sheets containing such fibers. The combination of carboxyethylated fibers and cationic additive materials has been found to be unexpectedly advantageous with regard to wet strength compared to combinations of carboxymethylated fibers and similar cationic additive materials. See U.S. Patent 25 No. 5,667,637.

Despite the advances in the use of carboxylated fibers and the formation of fibrous webs incorporating such fibers, there exists a need for carboxylated fibers that do not suffer the drawbacks of carboxymethylated and carboxyethylated cellulosic fibers, which include high cost and lost hemicelluloses. Accordingly, there is a need in 30 the art for modified cellulosic fibers having advantageous absorbent properties and, in

addition, having enhanced bondability so as to increase the strength of products that incorporate these fibers. The present invention seeks to fulfill these needs and offers further related advantages.

5

Summary of the Invention

In one aspect, the present invention provides carboxylated cellulosic fibers. Fibrous sheets and absorbent products containing carboxylated cellulosic fibers are also disclosed. The fibrous sheets generally include carboxylated fibers, a cationic additive, and, optionally, other fibers.

10

In another aspect of the invention, a method for producing carboxylated cellulosic fibers is provided. The method produces carboxylated cellulosic fibers by applying a carboxylating agent to the fibers and then heating the treated fibers for a period of time under controlled temperature, time, pH, and catalyst concentration conditions to effect bond formation between the carboxylating agent and the fiber while minimizing crosslinking reactions. The carboxylating agent is any chemical compound having two carboxylic acid groups separated by either two or three atoms such that the compound can form a cyclic 5- or 6-membered anhydride. Suitable carboxylating agents include succinic acid and succinic acid derivatives, phthalic acid, trimellitic acid, maleic acid, and itaconic acid and their derivatives. Bond formation between the carboxylating agent and the fiber is preferably the formation of a single ester bond between the carboxylating agent and the fiber and not the formation of extensive fiber crosslinks.

15

20

25

Brief Description of the Drawings

25

The foregoing aspects and many of the attendant advantages of this invention will become more readily appreciated by reference to the following detailed description, when taken in conjunction with the accompanying drawings, wherein:

30

FIGURE 1 is a graph showing wet burst strength of handsheets prepared from refined soft wood pulp (various Canadian Standard Freeness, CSF) modified with succinic acid (SUC) and 2 percent Kymene® 557H; GrP control refers to a handsheet

prepared from unmodified fibers; SUC-5.1 and SUC-7.1 refer to handsheets prepared from succinic acid-modified fibers having 5.1 and 7.1 milliequivalents (meq) carboxyl groups/100g fiber, respectively;

5 FIGURE 2 is a graph showing wet burst strength of handsheets prepared from refined soft wood pulp (various CSF) modified with sulfosuccinic acid (SULF) and 2 percent Kymene® 557H; GrP control refers to a handsheet prepared from unmodified fibers; SULF-7, SULF-13, and SULF-17 refer to handsheets prepared from sulfosuccinic acid-modified fibers having 7, 13, and 17 meq carboxyl groups/100g fiber, respectively;

10 FIGURE 3 is a graph showing wet burst strength of handsheets prepared from refined soft wood pulp (various CSF) modified with 2,2-dimethylsuccinic acid (DMS) and 2 percent Kymene® 557H; GrP control refers to a handsheet prepared from unmodified fibers; DMS-7, DMS-12, DMS-17, and DMS-25 refer to handsheets prepared from 2,2-dimethylsuccinic acid-modified fibers having 7, 12, 17, and 25 meq 15 carboxyl groups/100g fiber, respectively;

FIGURE 4 is a graph showing dry tensile strength of handsheets modified with 2,2-dimethylsuccinic acid (DMS) and 2 percent Kymene® 557H at various levels of refinement (CSF); GrP control refers to a handsheet prepared from unmodified fibers; DMS-7, DMS-12, DMS-17, and DMS-25 refer to handsheets prepared from 2,2-20 dimethylsuccinic acid-modified fibers having 7, 12, 17, and 25 meq carboxyl groups/100g fiber, respectively; and

FIGURE 5 is a graph showing the ratio of wet burst to dry tensile strength for handsheets modified with 2,2-dimethylsuccinic (DMS) and 2 percent Kymene® 557H at various levels of refinement (CSF); GrP control refers to a handsheet prepared from 25 unmodified fibers; DMS-7, DMS-12, DMS-17, and DMS-25 refer to handsheets prepared from 2,2-dimethylsuccinic acid-modified fibers having 7, 12, 17, and 25 meq carboxyl groups/100g fiber, respectively.

Detailed Description of the Preferred Embodiment

The present invention is directed to cellulosic fibers having enhanced bondability and methods related to such fibers. More specifically, the invention relates to carboxylated cellulosic fibers, products containing these cellulosic fibers, and 5 methods for producing and using these fibers. The carboxylated cellulosic fibers of the invention exhibit high absorbent capacity and bulk, and when such fibers are formed into a sheet and/or incorporated into an absorbent product, the resulting sheet or absorbent product exhibits increased wet strength in the presence of a cationic wet strength additive. The carboxylated cellulosic fibers of the invention can also be 10 advantageously combined with other fibers to provide a fibrous mixture having increased sheet strength.

In one aspect, the present invention provides a carboxylated cellulosic fiber having enhanced bondability and absorbent capacity. As used herein, the term "carboxylated cellulosic fiber" refers to a cellulosic fiber that has been modified to 15 include carboxylic acid groups (i.e., carboxyl groups) by chemical reaction with a carboxylating agent.

The carboxylating agent useful in forming the carboxylated cellulosic fiber of the invention is a chemical compound having two carboxylic acid groups separated by either two or three atoms such that the compound can form a cyclic 5- or 6-membered 20 anhydride ring. Generally, the carboxylating agent is a polycarboxylic acid. As used herein, the term "polycarboxylic acid" refers to an organic acid that contains two or more carboxylic acid groups, or the functional equivalent of two or more carboxylic acid groups, for example, acid salt, ester, and anhydride groups, among others. The carboxylated fiber includes a polycarboxylic acid covalently coupled or bonded to the 25 cellulose fiber. The polycarboxylic acid is coupled to the fiber through the formation of an ester bond between a carboxylic acid group on the polycarboxylic acid and a hydroxyl group on the cellulosic fiber. Coupling the polycarboxylic acid to the fiber in this way provides a fiber into which a carboxylic acid group has been incorporated. Where the carboxylating agent is a polycarboxylic acid having two carboxylic groups 30 (i.e., a dicarboxylic acid), the modified fiber preferably includes one carboxyl group for

each carboxylating agent reacted with and coupled to the fiber (i.e., the carboxylating agent provides one carboxyl equivalent to the fiber). For carboxylating agents that are polycarboxylic acids that contain three or more carboxylic acid groups, the modified fiber preferably includes more than one carboxyl group for each carboxylating agent 5 coupled to the fiber.

The carboxylated fibers of the present invention can vary with regard to the extent of incorporated carboxyl groups. Generally, sufficient carboxyl groups are incorporated into the fibers to provide an improvement in wet strength when combined with wet strength additives, absorbent capacity, or other advantageous property 10 compared to unmodified fibers. Depending on the nature of the subsequent use of a particular carboxylated fiber, the carboxylated fibers have from about 5 to about 50 milliequivalent (meq) carboxyl groups per 100 grams fiber. In a preferred embodiment, the carboxylated fibers have from about 6 to about 40 meq carboxyl groups per 100 grams fiber.

15 As noted above, the carboxylated fibers of this invention are produced by treating cellulosic fibers with a carboxylating agent, and optionally a catalyst, for a period of time and at a temperature sufficient to form an ester bond between the polycarboxylic acid and the fiber. In contrast to "curing", which refers to the exhaustive reaction of an agent (e.g., a crosslinking agent) with fibers, the bonding of 20 the polycarboxylic acid to the fibers in accordance with the present invention refers to less than exhaustive reaction of the polycarboxylic acid's carboxyl groups with the fiber. For example, for many crosslinking agents, including polycarboxylic acid crosslinking agents, exhaustive reaction between the fiber and substantially all of the crosslinking agent's carboxylic acid groups is desired and accomplished by either 25 prolonged reaction time and/or elevated cure temperature. Polycarboxylic acid "covalent coupling" or "bonding" to the fibers in accordance with the present invention refers to a controlled, nonexhaustive reaction, for example, the coupling of less than all carboxyl groups, and more preferably only a single carboxyl group, of the polycarboxylic acid to a fiber. An important aspect of the present invention is the 30 discovery of a method to accomplish coupling while minimizing or eliminating

crosslinking. Crosslinking reduces the interfiber bonding of fibers by reducing the swelling and water retention value (WRV) of wet fibers. Reduction of these properties results in reduced bonded area between fibers. Thus, a preferred embodiment of this invention includes conducting the coupling reaction such that the carboxylated fibers 5 have a WRV equal to that of the starting fibers, and preferably greater than that of the starting fibers.

Generally, the carboxylating agent useful in forming the carboxylated fibers of the invention is an organic acid containing two or more carboxyl groups having either a 1,2- or a 1,3-diacid substitution. That is, the carboxylating agent contains at least 10 two carboxylic acid groups with one carboxyl group separated from the second carboxyl group by either two (i.e., 1,2-diacid) or three (i.e., 1,3-diacid) atoms. Without being bound by theory, it appears that a carboxyl group is most reactive toward bonding with cellulose when the carboxylating agent can form a cyclic five- or 15 six-membered anhydride with a neighboring carboxyl group. Thus, the carboxylating agent useful in the present invention preferably contains at least two carboxyl groups that are separated by either two or three atoms in the chain or ring to which the carboxyl groups are attached. The atoms separating the carboxyl groups can include carbon, nitrogen, sulfur, and oxygen atoms, and mixture of these atoms. Preferably, the carboxylating agent includes two carboxyl groups that are separated by carbon 20 atoms, more preferably saturated carbon atoms (e.g., methylene and methine carbons) and carbon atoms that are further substituted (e.g., dimethyl and sulfonic acid substituted carbons).

Suitable carboxylating agents include aliphatic, unsaturated, aromatic, alicyclic and cyclic acids. For carboxylating agents having two carboxyl groups separated by a 25 carbon-carbon double bond (e.g., unsaturated acids) or where both carboxyl groups are connected to the same ring (e.g., cycloalkyl), the two carboxyl groups must be in a cis configuration relative to each other so that the carboxylating agent can form a cyclic five- or six-membered anhydride.

In a preferred embodiment, the carboxylating agent is a dicarboxylic acid 30 having two or three atoms separating the carboxyl groups. In one preferred

embodiment, the carboxylating agent is a 1,2-dicarboxylic acid or derivative, preferably succinic acid (i.e., HO₂CCH₂CH₂CO₂H) or a succinic acid derivative. Preferred succinic acid derivatives include 2-sulfosuccinic acid and 2,2-dimethylsuccinic acid. In another preferred embodiment, the carboxylating agent is a 5 1,3-dicarboxyl acid, preferably glutaric acid (i.e., HO₂CCH₂CH₂CH₂CO₂H) or a glutaric acid derivative. Preferred glutaric acid derivatives include 2,2-dimethylglutaric acid and diglycolic acid (i.e., HO₂CCH₂OCH₂CO₂H). Other suitable dicarboxylic acids include 1,2-dicarboxybenzene (e.g., 1,2-phthalic acid) and its derivatives, 1,2- and 1,3-dicarboxycycloalkanes, trimellitic acid, maleic acid, and itaconic acid and their 10 derivatives.

In the practice of the present invention, dicarboxylic acids having either a 1,2- or a 1,3-diacid substitution are preferred because the diacid can (1) form a cyclic five- or six-member anhydride, which is reactive toward cellulosic hydroxyl groups, and (2) provide a free carboxyl group that is relatively resistant to subsequent ester formation 15 with a cellulosic hydroxyl group. For the reasons noted above, the free carboxyl group incorporated into the fiber by carboxylating with a 1,2- or 1,3-dicarboxylic acid, or acid derivative, is resistant to subsequent ester formation with the cellulose fiber (i.e., the dicarboxylic acid does not function as a crosslinking agent). Preferred carboxylating agents ultimately form a single ester bond with a cellulose fiber and 20 incorporate one or more carboxyl groups for each carboxylating agent coupled to the fiber.

Polycarboxylic acids having more than two carboxyl groups have been previously utilized to effectively crosslink cellulose to provide cellulosic fibers having high bulk, resilience, and rapid liquid acquisition properties. Such crosslinked fibers 25 suffer from low bondability by virtue of the loss of interfiber hydrogen bonding that accompanies crosslinking. Basically, crosslinking reduces the relative bonded area between fibers by reducing swelling, conformability, flexibility, and surface area of wet fibers. Crosslinking also reduces the refinability of fibers, that is, the ability to create additional surface area through mechanical refining. Thus, although sheets of

crosslinked fibers have high bulk and certain advantageous absorbent properties, these sheets suffer from low dry and wet strength.

Despite the inherent disadvantages noted above associated with crosslinking cellulosic fibers with polycarboxylic acids, under certain conditions, polycarboxylic acids having three or more carboxy groups can be used in forming the carboxylated fibers of the present invention. When polycarboxylic acids are used as carboxylating agents, conditions for coupling the polycarboxylic acid to the fiber are such that exhaustive reaction (i.e., extensive crosslinking) is avoided and the polycarboxylic acid is preferably coupled to the fiber through a single ester bond and the remaining polycarboxylic acid's carboxyl groups are incorporated as free carboxyl groups to the fiber. Reaction conditions such as temperature, pH, time, fiber moisture content, crosslinking agent concentration, and catalyst concentration, among others, can be optimized to promote coupling of a polycarboxylic acid to fibers without significant crosslinking to provide carboxylated fibers having the advantageous properties noted above.

The carboxylated cellulosic fibers formed in accordance with the present invention include a polycarboxylic acid covalently coupled to a cellulose fiber through an ester bond. Although the polycarboxylic acid useful in the present invention is not a crosslinking agent, it will be appreciated that, while the formation of multiple ester bonds between a polycarboxylic acid and one or more cellulose chains or fibers is minimized, it can still occur to a limited extent and, therefore, such bonding between the polycarboxylic acid and the fibers is within the scope of this invention. For example, the polycarboxylic acid may form a single ester bond to a cellulose chain, two or more ester bonds with a chain, or two or more ester bonds between two or more chains or fibers. In any event, in accordance with the present invention, after covalent coupling to the fiber, the polycarboxylic acid has at least one free carboxylic acid group.

In addition to the dicarboxylic acids described above, other suitable carboxylating agents include polycarboxylic acids containing three or more carboxyl groups. Exemplary polycarboxylic acids include citric acid (i.e., 2-hydroxy-1,2,3-

propane tricarboxylic acid), 1,2,3-propane tricarboxylic acid, 1,2,3,4-butane tetracarboxylic acid, tartrate monosuccinic acid, tartrate disuccinic acid, oxydisuccinic acid (i.e., 2,2'-oxybis(butanedioic acid)), thiidisuccinic acid, trans-1-propene-1,2,3-tricarboxylic acid, all cis-1,2,3,4-cyclopentanetetracarboxylic acid, and 5 benzenehexacarboxylic acid.

In addition to the polycarboxylic acids described and noted above, polycarboxylic acid carboxylating agents include polymeric polycarboxylic acids. Suitable polymeric polycarboxylic acids include homopolymeric and copolymeric polycarboxylic acids and may advantageously incorporate self-catalyzing substituents 10 in the polymer chain, such as phosphonoalkyl groups. Representative homopolymeric polycarboxylic acids include, for example, polyacrylic acid, polyitaconic acid, and polymaleic acid. Examples of representative copolymeric polycarboxylic acids include polyacrylic acid copolymers such as poly(acrylamide-co-acrylic acid), poly(acrylic acid-co-maleic acid), poly(ethylene-co-acrylic acid), and poly(1-vinylpyrrolidone-co- 15 acrylic acid), as well as other polycarboxylic acid copolymers including poly(ethylene-co-methacrylic acid), poly(methyl methacrylate-co-methacrylic acid), poly(methyl vinyl ether-co-maleic acid), poly(styrene-co-maleic acid), and poly(vinyl chloride-co-vinyl acetate-co-maleic acid). In one preferred embodiment, the polymeric polycarboxylic acid is a polyacrylic acid. In another preferred embodiment, the polycarboxylic acid is 20 a polyacrylic acid containing phosphonoalkyl groups (e.g., A9930 commercially available from Rohm and Haas, Co., Philadelphia, PA). In another preferred embodiment, the polymeric polycarboxylic acid is a polymaleic acid. In still another preferred embodiment, the polymeric polycarboxylic acid is copolymer of acrylic acid, and preferably a copolymer of acrylic acid and another acid, for example, maleic acid. 25 The representative polycarboxylic acids noted above are available in various molecular weights and ranges of molecular weights from commercial sources.

In contrast to the polyacrylic acid crosslinking agent treatment described in Herron, in the method of the present invention the polycarboxylic acids are not subjected to elevated cure temperatures to effect exhaustive polycarboxylic acid-to- 30 fiber crosslinking. Rather, in this invention, the polycarboxylic acid is cured at a

significantly lower temperature to accomplish the opposite effect, namely, to effect covalent coupling of the carboxylic acid to the fibers and at the same time, maintain sufficient free carboxylic acid groups (i.e., carboxylic acid groups that are not bonded to the fiber) to impart the advantageous properties of absorbent capacity and bondability to the fibers, and absorbency and strength to fibrous compositions incorporating these fibers. In the context of the present invention, the polycarboxylic acid is optimally covalently coupled to the fiber through a single carboxylic acid group, forming a single ester bond between the fiber and the polycarboxylic acid. Reaction through a single carboxylic acid group allows the remaining carboxylic acid group or groups of the polycarboxylic acid to participate in interfiber interactions (e.g., hydrogen bonding) in fibrous compositions, thereby enhancing the strength of those compositions. Thus, although the invention described in the Herron patents and the present invention generally incorporate a polycarboxylic acid into cellulose fibers, because of the diverse treatments and goals, the resulting products are distinct. As noted above, the Herron patents describe utilizing a polycarboxylic acid as a crosslinking agent to form intrafiber ester crosslinks. In contrast, the present invention utilizes a polycarboxylic acid as a carboxylating agent to incorporate one or more carboxyl groups into the fiber to enhance the fibers' bondability.

Those knowledgeable in the area of polycarboxylic acids will recognize that the polycarboxylic acids useful in the present invention may be present on the fibers in a variety of forms including, for example, the free acid form, and salts thereof. It will be appreciated that all such forms are included within the scope of the invention. Furthermore, although the carboxylating agent has been described as a polycarboxylic acid, it will be appreciated that other carboxylating agents that include functional groups capable of providing a polycarboxylic acid, for example, an acid salt, an ester, or an acid anhydride, having the properties and characteristics described above are also carboxylating agents within the scope of this invention.

The carboxylating agents noted above can be used alone or in combination to provide the cellulose fibers of the present invention having carboxyl groups.

The carboxylated cellulose fibers have an effective amount of a polycarboxylic acid covalently coupled to the fibers through an ester bond. That is, polycarboxylic acid in an amount sufficient to provide an improvement in strength (e.g., tensile, sheet) in compositions (e.g., fibrous sheets, webs, mats) containing the cellulose fibers to 5 which the polycarboxylic acid is covalently coupled, relative to conventional fibers lacking such carboxylated fibers. Generally, the cellulose fibers are treated with a sufficient amount of a polycarboxylic acid such that an effective amount of polycarboxylic acid is covalently coupled to the fibers.

The polycarboxylic acid is preferably present on the fibers in an amount from 10 about 0.1 to about 10 percent by weight of the total weight of the fibers. More preferably, the polycarboxylic acid is present in an amount from about 0.2 to about 7 percent by weight of the total weight of the fibers, and in a particularly preferred embodiment, from about 0.4 to about 6 percent by weight of the total weight of the fibers. At less than about 0.1 percent by weight polycarboxylic acid, no significant 15 absorbent or bondability enhancement is observed, and at greater than about 10 percent by weight, the maximum coupling capacity of the fibers is exceeded.

The carboxylating agent can be applied to the fibers for covalent coupling by any one of a number of methods known in the production of treated fibers. For example, the carboxylating agent can be contacted with the fibers as a fiber sheet is 20 passed through a bath containing the carboxylating agent. Alternatively, other methods of applying the carboxylating agent, including fiber spraying, or spraying and pressing, or dipping and pressing with a carboxylating agent solution, are also within the scope of the invention.

Generally, the carboxylated cellulosic fibers of the present invention can be 25 prepared by applying a carboxylating agent, as described above, to cellulose fibers, and then coupling or bonding the carboxylating agent to the fibers for a period of time and at a temperature sufficient to effect ester bond formation between the carboxylating agent and the fibers. In the context of the present invention, such ester bond formation between the carboxylating agent and fibers is not exhaustive ester bond formation as in 30 fiber crosslinking. The temperature sufficient to effect ester bond formation is

generally lower than the cure temperature of a typical crosslinking agent and will also vary depending upon the specific acid and moisture content of the fibers, among other factors. For an exemplary acid, succinic acid, the temperature sufficient to effect ester bond formation ranges from about 120°C to about 160°C. The use of a catalyst to 5 promote ester bond formation between the carboxylating agent and the cellulose fiber in the method is preferred and reduces the temperature required to effect ester bond formation. While catalysts can be used to effectively lower the bonding temperature of the carboxylating agent, in accordance with the present invention, the use of catalysts preferably does not result in exhaustive crosslinking of the carboxylating agent to the 10 fibers. The effect of bonding temperature on the introduction of carboxylic acid groups and water retention value for fibers treated with succinic acid is summarized in Example 1, Table 1. It can be seen that the WRV maximum is at 130°C to 140°C and that at higher bonding temperatures the WRV decreases due to a higher proportion of crosslinking reactions.

15 As noted above, the carboxylated cellulosic fibers of the invention can also be prepared with the aid of a catalyst. In such a method, the catalyst is applied to the cellulose fibers in a manner analogous to application of the carboxylating agent to the fibers as described above. The catalyst may be applied to the fibers prior to, after, or at the same time that the carboxylating agent is applied to the fibers. Accordingly, the 20 present invention provides a method of producing carboxylated cellulosic fibers that includes coupling the carboxylating agent to the fibers in the presence or absence of a catalyst.

Generally, the catalyst promotes ester bond formation between the carboxylating agent and the cellulose fibers and is effective in increasing bond 25 formation (i.e., the number of bonds formed) at a given cure temperature. Suitable catalysts include any catalyst that increases the rate of bond formation between the carboxylating agent and cellulose fibers. Preferred catalysts include alkali metal salts of phosphorous containing acids such as alkali metal hypophosphites, alkali metal phosphites, alkali metal polyphosphonates, alkali metal phosphates, and alkali metal sulfonates. Particularly preferred catalysts include alkali metal polyphosphonates such 30

as sodium hexametaphosphate, and alkali metal hypophosphites such as sodium hypophosphite. When a catalyst is used to promote bond formation, the catalyst is typically present in an amount in the range from about 5 to about 20 weight percent of the carboxylating agent. Preferably, the catalyst is present in about 10 percent by 5 weight of the carboxylating agent. The effect of catalyst (1.5 to 3.0 percent by weight sodium hypophosphite at 140°C) on the introduction of carboxylic acid groups and water retention value for fibers treated with succinic acid is summarized in Example 1, Table 2.

Cellulosic fibers are a basic component of the carboxylated fibers of the present 10 invention. Although available from other sources, cellulosic fibers are derived primarily from wood pulp. Suitable wood pulp fibers for use with the invention can be obtained from well-known chemical processes, such as the kraft and sulfite processes, with or without subsequent bleaching. The pulp fibers may also be processed by thermomechanical, chemithermomechanical methods, or combinations thereof. The 15 preferred pulp fiber is produced by chemical methods. Ground wood fibers, recycled or secondary wood pulp fibers, and bleached and unbleached wood pulp fibers can be used. Softwoods and hardwoods can be used. Details of the selection of wood pulp fibers are well-known to those skilled in the art. These fibers are commercially available from a number of companies, including Weyerhaeuser Company, the assignee 20 of the present invention. For example, suitable cellulose fibers produced from southern pine that are usable with the present invention are available from Weyerhaeuser Company under the designations CF416, NF405, PL416, FR516, and NB416.

In general, the carboxylated cellulosic fibers of the present invention may be 25 prepared by a system and apparatus as described in U.S. Patent No. 5,447,977 to Young, Sr. et al., which is incorporated herein by reference in its entirety. Briefly, the fibers are prepared by a system and apparatus comprising a conveying device for transporting a mat of cellulose fibers through a fiber treatment zone; an applicator for applying a treatment substance such as a carboxylating agent to the fibers at the fiber 30 treatment zone; a fiberizer for completely separating the individual cellulosic fibers

comprising the mat to form a fiber output comprised of substantially unbroken and individualized cellulose fibers; and a dryer coupled to the fiberizer for flash evaporating residual moisture and for bonding the carboxylating agent to the fiber and to form dried, individualized carboxylated fibers.

5 As used herein, the term "mat" refers to any nonwoven sheet structure comprising cellulose fibers or other fibers that are not covalently bound together. As noted above, fibers include those obtained from wood pulp or other sources including cotton rag, hemp, grasses, cane, husks, cornstalks, or other suitable sources of cellulose fibers that can be laid into a sheet. The mat of cellulose fibers is preferably in
10 an extended sheet form, and can be one of a number of baled sheets of discrete size or can be a continuous roll.

Each mat of cellulose fibers is transported by a conveying device, for example, a conveyor belt or a series of driven rollers. The conveying device carries the mats through the fiber treatment zone.

15 At the fiber treatment zone the carboxylating agent acid is applied to the cellulose fibers. The carboxylating agent is preferably applied to one or both surfaces of the mat using any one of a variety of methods known in the art including spraying, rolling, or dipping. Once the materials have been applied to the mat, the materials can be uniformly distributed through the mat, for example, by passing the mat through a
20 pair of rollers.

After the fibers have been treated with the carboxylating agent, the impregnated mat can be fiberized by feeding the mat through a hammermill. The hammermill serves to separate the mat into its component individual cellulose fibers, which are then blown into a dryer.

25 The dryer performs two sequential functions; first removing residual moisture from the fibers, and second bonding the carboxylating agent in accordance with the present invention. In one embodiment, the dryer comprises a first drying zone for receiving the fibers and for removing residual moisture from the fibers via a flash-drying method, and a second drying zone for effecting the carboxylating agent-to-fiber
30 bond. Alternatively, in another embodiment, the treated fibers are blown through a

flash-dryer to remove residual moisture, and then transferred to an oven where the treated fibers are subsequently formed in accordance with the present invention.

A representative method for forming the carboxylated fibers of the invention is described in Example 1. The incorporation of carboxylic acid groups and water 5 retention values for representative carboxylated fibers prepared by treating with succinic acid are presented in Example 1, Tables 1-3. As noted above, the present invention provides carboxylated fibers having a water retention value about equal to, preferably greater than, the water retention value of fibers from which the carboxylated fibers are formed. In general, the carboxylated fibers of the invention have a water 10 retention value greater than about 1.0 g/g. Generally, increasing carboxylic acid group incorporation into the fibers increases the fibers' water retention value. However, at higher bonding temperatures, increased carboxylic acid group incorporation can be accompanied by increased crosslinking, which results in a decrease in the fibers' water retention value. Increased incorporation of carboxylic acid groups into the fibers also 15 increases the fibers' bondability. In a preferred method, fibers are treated with a carboxylating agent (about 6 percent by weight based on total weight of fibers) at pH of from about 2 to about 4 in the presence of a catalyst (about 3 percent by weight based on total weight of fibers) and then heated at about 140°C to effect carboxylating agent-to-fiber bonding.

20 The carboxylated cellulosic fibers of the present invention are preferably combined with a cationic additive to form fibrous sheets and absorbent products that exhibit enhanced wet and/or dry strength. The advantageous strength properties imparted to fibrous compositions that include carboxylated fibers and a cationic additive are due, at least in part, to the relatively strong attraction and association of 25 the cationic additive to the carboxylated fibers, which are anionic in nature.

Exemplary cationic additives include, for example, wet strength resins and cationic starches that are useful in paper manufacturing. Suitable wet strength resins include polyamide epichlorohydrin, polyethyleneimine, and polyacrylamide wet strength resins. Polyamide epichlorohydrin resin is commercially available, for 30 example, under the designation Kymene® 557LX and 557H (Hercules, Inc.,

Wilmington, DE). Polyacrylamide resin is described, for example, in U.S. Patent No. 3,556,932 issued January 19, 1971 to Coscia et al., and another is commercially available under the designation ParezTM 631 NC (American Cyanamid Co., Stamford, CT). Cationic starches are commercially available from a variety of sources including

5 National Starch and Chemical Corp., Bridgewater, NJ. A preferred cationic starch is available from Western Polymer Co., Moses Lake, WA under the designation Wescat EF. A general discussion on wet strength resins utilized in the paper field, and generally applicable in the present invention, can be found in TAPPI Monograph Series No. 29, "Wet Strength in Paper and Paperboard", Technical Association of the Pulp

10 and Paper Industry (New York, 1965), expressly incorporated herein in its entirety.

Generally, the wet strength agent is present in the composition in an amount from about 0.01 to about 10 weight percent, and preferably from about 0.1 to about 5 weight percent, based on the total weight of the composite. In one preferred embodiment, the wet strength agent useful in forming the composite of the present

15 invention is a polyamide epichlorohydrin resin commercially available from Hercules, Inc. under the designation Kymene[®] 557H. The wet and dry tensile strengths of an absorbent composite formed in accordance with the present invention will generally increase with an increase in the amount of wet strength agent.

Carboxylated fibers that further include a cationic additive can also be prepared

20 as generally described above. Briefly, such fibers can be prepared by applying a cationic additive to the fibrous mat, for example, at the fiber treatment zone. The cationic additive can be applied to the fibrous mat either before, during, or after application of the carboxylating agent. The resulting treated fibers can then be fiberized and heated to effect drying and bonding of the carboxylating agent to the

25 fibers to provide individualized carboxylated fibers that further include a cationic additive.

Alternatively, a fibrous mat or web can be formed by applying a carboxylating agent and, optionally, a cationic additive, to the fibrous mat and, rather than fiberizing the mat to form individualized fibers, the treated fibrous mat can be heated to effect

30 drying and bonding of the carboxylating agent to the fibers to provide a mat of

carboxylated fibers. Such a mat is particularly useful for transporting carboxylated fibers to subsequent destinations where the mat can then be fiberized to provide individual fibers that can be further combined with other fibers and materials as desired to provide various absorbent products. The carboxylated fibrous mat further including 5 a cationic additive can also be subsequently reslurried and combined with other fibers and materials to provide a variety of fibrous products.

The carboxylated cellulosic fibers formed as described above are fibers that have been modified to include carboxyl groups. The modified fibers' carboxyl groups are available to form hydrogen bonds with, for example, other fibers including other 10 carboxylated fibers. Therefore, the carboxylated fibers formed in accordance with the present invention, optionally including a cationic additive, can be advantageously combined with other fibers and materials to provide a fibrous composite having a variety of properties including advantageous strength properties imparted to the composite by the carboxylated fibers. The carboxylated fibers of the invention, 15 optionally including a cationic additive, can be combined with other fibers including carboxylated fibers such as carboxymethylcellulose and carboxyethylcellulose, crosslinked cellulosic fibers, untreated cellulosic fibers, thermomechanical fibers, chemithermomechanical (CTMP) fibers, cellulose acetate fibers, polyester fibers, and thermobondable fibers.

20 A representative procedure for forming fibrous webs that include the carboxylated fibers of the invention is described in Example 2. Generally, fibrous webs formed from carboxylated fibers and a wet strength agent have increased wet strength compared to fibrous webs that do not contain carboxylated fibers. The wet burst strength of handsheets formed from carboxylated fibers and a representative wet 25 strength agent was found to be significantly greater than for handsheets prepared from the corresponding untreated fibers. FIGURES 1-3 illustrate the increase in wet burst strength for handsheets formed from fibers treated with 2 percent Kymene® 557H and various amounts of succinic acid, sulfosuccinic acid, and 2,2-dimethylsuccinic acid, respectively.

Fibrous webs formed from the carboxylated fibers of the invention also have reduced dry strength compared to webs formed from untreated fibers. Reduced web dry strength corresponds to enhanced web softness. Thus, incorporating carboxylated fibers into a fibrous web provides a web with enhanced softness compared to a 5 corresponding web prepared from untreated fibers. The dry tensile strength of representative handsheets formed from carboxylated (i.e., 2,2-dimethylsuccinic acid) fibers and a wet strength agent (i.e., 2 percent Kymene®) and a corresponding handsheet formed from untreated fibers is illustrated in FIGURE 4. Referring to FIGURE 4, the dry tensile strength of the handsheets formed from the carboxylated 10 fibers is significantly reduced compared to the web formed from untreated fibers. The ratio of wet burst strength to dry tensile strength for handsheets prepared from carboxylated fibers and containing a wet strength agent (i.e., 2 percent Kymene®) is illustrated in FIGURE 5. Referring to FIGURE 5, the high wet/dry strength ratio for the handsheets formed in accordance with the present invention compared to 15 handsheets formed from untreated fibers indicates that the handsheets that include carboxylated fibers possess advantageous wet strength in addition to softness.

Carboxylated cellulosic fibers provide advantageous absorbent and strength properties to fibrous composites that include such fibers. By virtue of bonding the carboxylating agent to the fiber, anionic sites and hydrogen bonding sites are added to 20 the fiber. Generally, the carboxyl groups enhance fiber swelling, which provides for advantageous absorbent properties. In addition, the carboxyl groups provide for strong attraction and association to cationic additives such as wet strength agents that increase the wet strength and integrity of absorbent products that include these fibers.

The carboxylated fibers of the invention can be formed into sheets or mats 25 having high absorbent capacity, bulk, resilience, and increased tensile strength. For example, these fibers may be combined with other fibers such as crosslinked and CTMP pulp fibers. The resulting sheets can be incorporated into a variety of absorbent products including, for example, tissue sheets, paper toweling, disposable diapers, adult incontinence products, sanitary napkins, and feminine care products.

The carboxylated fibers of the present invention are particularly useful in absorbent products requiring high wet burst strength.

The following examples illustrate the practice of the present invention, and are not intended to be limiting thereof.

5

EXAMPLES

Example 1

A Representative Method for Preparing Carboxylated Cellulosic Fibers

The carboxylated cellulosic fibers of the present invention and products 10 containing these fibers can be prepared by a system and apparatus as generally described in U.S. Patent No. 5,447,977 to Young, Sr. et al., which is incorporated herein by reference in its entirety.

In this example, the preparation of carboxylated cellulosic fibers is described. This example demonstrates that a polycarboxylic acid can be bonded to cellulosic 15 fibers to provide fibers having enhanced absorbent capacity and bondability.

In the process, a fiber sheet composed of individual cellulose fibers (available under the designation NB416 from Weyerhaeuser Co., New Bern, NC) is treated with succinic acid at varying bonding temperatures according to the following procedure.

Briefly, a fiber sheet is fed from a roll through a constantly replenished bath of 20 an aqueous solution containing succinic acid adjusted to concentrations to achieve the desired level of succinic acid (e.g., about 0.25 to about 10 percent by weight of the total composition) and sodium hypophosphite (at a concentration approximately one-half that of succinic acid). The treated fiber sheet is then moved through a roller nip set to remove sufficient solution to provide a fiber sheet having a pulp solids content of 25 about 50 percent. After passing through the roll nip, the wet fibrous sheet is air dried. The bonding of the polycarboxylic acid to the individualized fibers is completed by placing the fibrous sheet in a laboratory oven and heating at about 140°C for 20 minutes.

The effect of bonding temperature on the level of carboxylic acid group 30 incorporation into the fibers and the water retention value of the fibers is summarized

in Table 1. Fibers were treated with succinic acid (6 percent by weight based on the total weight of fibers) and sodium hypophosphite (3 percent by weight based on the total weight of fibers) and heated at the indicated temperature for 20 minutes. Water retention value (WRV) was determined by TAPPI Method UM 256, and the level of 5 carboxylic acid group incorporation was determined by TAPPI Method T237 OM-88. In Table 1, Control 120 and Control 160 refer to control fibers that were heated to the respective bonding temperature without succinic acid treatment. Yield (%) refers to the percent conversion of succinic acid.

10

Table 1. The Effect of Temperature on Succinic Acid Esterification of Cellulose Fibers

Temp. °C	Carboxyl Level (meq/100g)	WRV (g/g)	Yield (%)
120	12	1.22	25
130	23	1.31	46
140	26	1.31	53
150	30	1.29	60
160	34	0.96	67
Control 120	4	1.12	--
Control 160	4	1.00	--

The maximum WRV, and thus the maximum swelling of the fibers, is obtained at bonding temperatures of 130° to 140°C. Despite the fact that more carboxyl groups 15 are incorporated at higher temperatures, which would normally increase WRV and swelling, the WRV actually decreases due to the occurrence of undesirable crosslinking at temperatures above 140°C. The temperatures in Table 1 represent a 20-minute bonding time. As would be expected with any chemical reaction, the optimum temperature will increase with shorter bonding times, and decrease with 20 longer bonding times.

The effect of a catalyst on the bonding of the carboxylating agent to the fibers is summarized in Table 2. Fibers were treated with succinic acid (6 percent by weight based on total weight of fibers) and the indicated amount of sodium hypophosphite and heated at 140°C for 20 minutes.

Table 2. The Effect of Catalyst on Succinic Acid Esterification of Cellulose Fibers

Catalyst %	Carboxyl Level (meq/100g)	WRV (g/g)	Yield (%)
0	8	0.94	16
1.5	30	1.30	60
3.0	34	1.36	68

5 With no catalyst present, only a slight amount of esterification occurs, and the WRV of the fibers actually decreases instead of increasing. The result suggests that substantial crosslinking is occurring. With catalyst present in an effective amount, significantly more esterification occurs and the WRV of the fibers increases substantially.

10 The effect of pH on the bonding of the carboxylating agent to the fibers is summarized in Table 3. Fibers were treated with succinic acid (6 percent by weight based on the total weight of fibers) and sodium hypophosphite (3 percent by weight based on the total weight of fibers) and heated at 140°C for 20 minutes.

15 Table 3. The Effect of pH on Succinic Acid Esterification of Cellulose Fibers

pH	Carboxyl Level (meq/100g)	WRV (g/g)	Yield (%)
4.5	11	1.01	21
4.0	16	1.28	34
3.5	21	1.34	42
3.0	24	1.28	48
2.5	26	1.29	53
2.0	28	1.29	56

The effect of increasing the pH of the succinic acid/sodium hypophosphite solution from 2.0 up to 4.5 is to decrease the level of esterification proportionately. 20 However, the WRV and fiber swelling reach a maximum at pH 3.5. The results suggest that at pHs lower than 3.5, a higher degree of crosslinking occurs compared to pH 3.5 and above.

Example 2A Representative Method for Preparing Handsheets
Containing Carboxylated Cellulosic Fibers

5 In this example, the preparation of handsheets from representative carboxylated cellulosic fibers is described.

About 30.5 g of GrP pulp was refined in a PFI Refiner to the desired freeness as measured by the Canadian Standard Freeness (CSF) test. GrP (Grand Prairie Softwood) refers to a Canadian bleached kraft wood pulp made from a mixed furnish 10 predominantly of white spruce, lodgepole pine, and balsam fir, with the major component being spruce. The refiner was designated No. 138 manufactured by P.F.I. Mølle, Hamjern, Oslo, Norway. The freeness tester is manufactured by Robert Mitchell Company, Ltd., Ste. Laurent, Quebec. The refined pulp was then placed in a disintegrator for 10,000 revolutions to obtain a uniform slurry. The pulp slurry was 15 then diluted to 10 L and consistency determined. The disintegrator is a British Pulp Evaluation Apparatus, manufactured by Mavis Engineering, Ltd., London, England. All three machines are also available from Testing Machines Inc., Amityville, NY.

The cationic wet strength additive was a water-soluble polyamide epichlorohydrin (PAE) reaction product, Kymene® 557H (Hercules, Inc., Wilmington 20 DE). Kymene® 557H is supplied as a 12.5% solids aqueous solution. For use, Kymene® as received was diluted to a 1% solids solution.

Handsheets were formed in a conventional manner in a sheet mold that produced sheets 152 mm (6 in) in diameter. White water from the sheet mold was recycled as dilution water for subsequent sheets to better simulate commercial 25 operating conditions. The first seven sheets made were discarded to allow white water fines to build up to an equilibrium level. Following that, the eighth sheet was used to check sheet weight and adjust amount of stock added in order to produce the desired 1.2 g (oven dry weight) sheets. Then 10 additional sheets were made for testing.

Following drying, the sheets were oriented on edge in a wire rack and placed in 30 an oven at 100° C for one hour to allow good curing of any wet strength resin. A

number of samples were made using 100 percent modified carboxylated pulps as well as blends of these pulps with unmodified pulp. For most conditions, similar handsheet samples of the carboxylated pulps were made for comparison.

Physical properties of the various modified materials and blends are best 5 understood by referring to FIGURES 1-5. Wet burst tests were conducted using a Thwing-Albert Model 1300-177 Burst Tester (Thwing-Albert Instrument Co., Philadelphia, PA). Dry tensile tests were performed according to TAPPI Method 494 Tensile Breaking Properties of Paper and Paperboard.

While the preferred embodiment of the invention has been illustrated and 10 described, it will be appreciated that various changes can be made therein without departing from the spirit and scope of the invention.

The embodiments of the invention in which an exclusive property or privilege is claimed are defined as follows:

1. Carboxylated cellulosic fibers, comprising cellulosic fibers covalently coupled to a carboxylating agent through an ester bond, wherein the carboxylating agent provides a carboxyl group to the fibers, and wherein the carboxylating agent is a polycarboxylic acid having one carboxyl group separated from a second carboxyl group by either two or three atoms, wherein the carboxylated fibers have a water retention value greater than or equal to the water retention value of the fibers from which the carboxylated fibers are formed.
5
- 10 2. The fibers of Claim 1 wherein the carboxylated cellulosic fibers are individualized fibers.
3. The fibers of Claim 1 wherein the polycarboxylic acid comprises a 1,2-dicarboxylic acid.
- 15 4. The fibers of Claim 3 wherein the 1,2-dicarboxylic acid is selected from the group consisting of succinic acid, 2,2-dimethylsuccinic acid, 2-sulfosuccinic acid, maleic acid, their derivatives, and mixtures thereof.
5. The fibers of Claim 1 wherein the polycarboxylic acid comprises a 1,3-dicarboxylic acid.
- 20 6. The fibers of Claim 5 wherein the 1,3-dicarboxylic acid is selected from the group consisting of glutaric acid, 2,2-dimethylglutaric acid, diglycolic acid, their derivatives, and mixtures thereof.
7. The fibers of Claim 1 wherein the polycarboxylic acid comprises an organic acid having three or more carboxyl groups.

8. The fibers of Claim 7 wherein the polycarboxylic acid is selected from the group selected from citric acid, 1,2,3-tricarboxypropane, 1,2,3,4-tetracarboxybutane, their derivatives, and mixtures thereof.

9. The fibers of Claim 7 wherein the polycarboxylic acid comprises a 5 polymeric polycarboxylic acid.

10. The fibers of Claim 9 wherein the polymeric polycarboxylic acid is selected from the group consisting of polyacrylic acid, polymaleic acid, copolymers of acrylic acid, copolymers of maleic acid, a copolymer of acrylic and maleic acids, their derivatives, and mixtures thereof.

10 11. The fibers of Claim 1 wherein the polycarboxylic acid is coupled to the fiber through a single ester bond.

12. The fibers of Claim 1 wherein the cellulosic fiber is a wood pulp fiber.

13. The fibers of Claim 1 wherein the polycarboxylic acid is present on the fibers in an amount from about 0.1 to about 10 percent by weight of the fibers.

15 14. The fibers of Claim 1 having from about 5 to about 50 meq carboxyl groups per 100 grams of fiber.

15. A fibrous composition comprising carboxylated cellulosic fibers and a cationic additive, wherein the carboxylated cellulosic fibers comprise cellulosic fibers covalently coupled to a carboxylating agent through an ester bond, wherein the 20 carboxylating agent provides a carboxyl group to the fibers, and wherein the carboxylating agent is a polycarboxylic acid having one carboxyl group separated from a second carboxyl group by either two or three atoms, wherein the carboxylated fibers have a water retention value greater than or equal to the water retention value of the fibers from which the carboxylated fibers are formed.

16. The composition of Claim 15 wherein the polycarboxylic acid is selected from the group consisting of a dicarboxylic acid, an organic acid having three or more carboxyl groups, a polymeric polycarboxylic acid, and mixtures thereof.

17. The composition of Claim 16 wherein the dicarboxylic acid is selected 5 from the group consisting of succinic acid, 2,2-dimethylsuccinic acid, 2-sulfosuccinic acid, glutaric acid, 2,2-dimethylglutaric acid, diglycolic acid, their derivatives, and mixtures thereof.

18. The composition of Claim 15 wherein the cationic additive is selected from the group consisting of cationic starches and wet strength resins.

10 19. The composition of Claim 18 wherein the wet strength resin is selected from the group consisting of polyamide epichlorohydrin resins, polyethyleneimine resins, and polyacrylamide resins.

20. The composition of Claim 15 wherein the cationic additive comprises a polyamide epichlorohydrin resin.

15 21. The composition of Claim 15 wherein the cationic additive is present in about 0.01 to about 10 percent by weight based on the total weight of the composition.

20 22. A fibrous sheet comprising carboxylated cellulosic fibers, the fibers comprising cellulosic fibers covalently coupled to a carboxylating agent through an ester bond, wherein the carboxylating agent provides a carboxyl group to the fibers, and wherein the carboxylating agent is a polycarboxylic acid having one carboxyl group separated from a second carboxyl group by either two or three atoms, wherein the carboxylated fibers have a water retention value greater than or equal to the water retention value of the fibers from which the carboxylated fibers are formed.

23. The sheet of Claim 22 wherein the polycarboxylic acid is selected from the group consisting of a dicarboxylic acid, an organic acid having three or more carboxyl groups, a polymeric polycarboxylic acid, and mixtures thereof.

24. The sheet of Claim 22 wherein the dicarboxylic acid is selected from the 5 group consisting of succinic acid, 2,2-dimethylsuccinic acid, 2-sulfosuccinic acid, glutaric acid, 2,2-dimethylglutaric acid, diglycolic acid, their derivatives, and mixtures thereof.

25. The sheet of Claim 22 further comprising a cationic additive.

26. The sheet of Claim 22 wherein the cationic additive comprises a 10 polyamide epichlorohydrin resin.

27. The sheet of Claim 22 further comprising fibers selected from the group consisting of carboxymethylated fibers, carboxyethylated fibers, crosslinked fibers, untreated cellulosic fibers, thermomechanical fibers, chemithermomechanical fibers, cellulose acetate fibers, polyester fibers, thermobondable fibers, and mixtures thereof.

15 28. The sheet of Claim 25 further comprising fibers selected from the group consisting of carboxymethylated fibers, carboxyethylated fibers, crosslinked fibers, untreated cellulosic fibers, thermomechanical fibers, chemithermomechanical fibers, cellulose acetate fibers, polyester fibers, thermobondable fibers, and mixtures thereof.

29. A method for preparing individualized, carboxylated cellulosic fibers, 20 comprising:

applying a carboxylating agent to a fibrous mass, wherein the carboxylating agent is a polycarboxylic acid having one carboxyl group separated from a second carboxyl group by either two or three atoms;

25 separating the fibrous mass into individual, substantially unbroken fibers; and heating the individualized fibers to form an ester bond between the carboxylating agent and the fibers, wherein the carboxylated fibers have a water

retention value greater than or equal to the water retention value of the fibers from which the carboxylated fibers are formed.

30. The method of Claim 29 wherein the polycarboxylic acid is selected from the group consisting of a dicarboxylic acid, an organic acid having three or more carboxyl groups, a polymeric polycarboxylic acid, and mixtures thereof.

31. The method of Claim 29 further comprising applying a catalyst to the fibrous mass.

32. The method of Claim 31 wherein the catalyst comprises sodium hypophosphite.

10 33. The method of Claim 29 further comprising applying a cationic additive to the fibrous mass.

34. The method of Claim 33 wherein the cationic additive comprises a polyamide epichlorohydrin resin.

15 35. A method for preparing a carboxylated cellulosic fibrous web, comprising:

applying a carboxylating agent to a fibrous web, wherein the carboxylating agent is a polycarboxylic acid one carboxyl group separated from a second carboxyl group by either two or three atoms; and

20 heating the fibers to form an ester bond between the carboxylating agent and the fibers, wherein the carboxylated fibers have a water retention value greater than or equal to the water retention value of the fibers from which the carboxylated fibers are formed.

25 36. The method of Claim 35 wherein the polycarboxylic acid is selected from the group consisting of a dicarboxylic acid, an organic acid having three or more carboxyl groups, a polymeric polycarboxylic acid, and mixtures thereof.

37. The method of Claim 35 further comprising applying a catalyst to the fibrous mass.

38. The method of Claim 37 wherein the catalyst comprises sodium hypophosphite.

5 39. The method of Claim 35 further comprising applying a cationic additive to the fibrous mass.

40. The method of Claim 39 wherein the cationic additive comprises a polyamide epichlorohydrin resin.

41. A method for increasing the wet strength of a fibrous sheet, comprising:
10 forming a fibrous slurry, wherein the slurry comprises carboxylated cellulosic fibers, the fibers comprising cellulosic fibers covalently coupled to a carboxylating agent through an ester bond, wherein the carboxylating agent provides a carboxyl group to the fibers, and wherein the carboxylating agent is a polycarboxylic acid having one carboxyl group separated from a second carboxyl group by either two or
15 three atoms, wherein the carboxylated fibers have a water retention value greater than or equal to the water retention value of the fibers from which the carboxylated fibers are formed;

20 depositing the fibrous slurry on a foraminous support;
dewatering the deposited slurry to provide a wet composite; and
drying the wet composite to provide a fibrous sheet having a wet strength greater than a fibrous sheet formed from a fibrous slurry lacking carboxylated cellulosic fibers.

42. The method of Claim 41 wherein the polycarboxylic acid is selected from the group consisting of a dicarboxylic acid, an organic acid having three or more
25 carboxyl groups, a polymeric polycarboxylic acid, and mixtures thereof.

43. The method of Claim 41 wherein the fibrous slurry further comprises a catalyst.

44. The method of Claim 41 wherein the fibrous slurry further comprises a cationic additive.

45. The method of Claim 41 wherein the fibrous slurry comprises fibers selected from the group consisting of carboxymethylated fibers, carboxyethylated fibers, crosslinked fibers, untreated cellulosic fibers, thermomechanical fibers, chemithermomechanical fibers, cellulose acetate fibers, polyester fibers, thermobondable fibers, and mixtures thereof.

46. A method for enhancing the softness of a fibrous web, comprising forming a fibrous slurry, wherein the slurry comprises carboxylated cellulosic fibers, the fibers comprising cellulosic fibers covalently coupled to a carboxylating agent through an ester bond, wherein the carboxylating agent provides a carboxyl group to the fibers, and wherein the carboxylating agent is a polycarboxylic acid having one carboxyl group separated from a second carboxyl group by either two or three atoms, wherein the carboxylated fibers have a water retention value greater than or equal to the water retention value of the fibers from which the carboxylated fibers are formed;

depositing the fibrous slurry on a foraminous support;

dewatering the deposited slurry to provide a wet composite; and

drying the wet composite to provide a fibrous web having a softness greater than a fibrous web formed from a fibrous slurry lacking carboxylated cellulosic fibers.

1/5

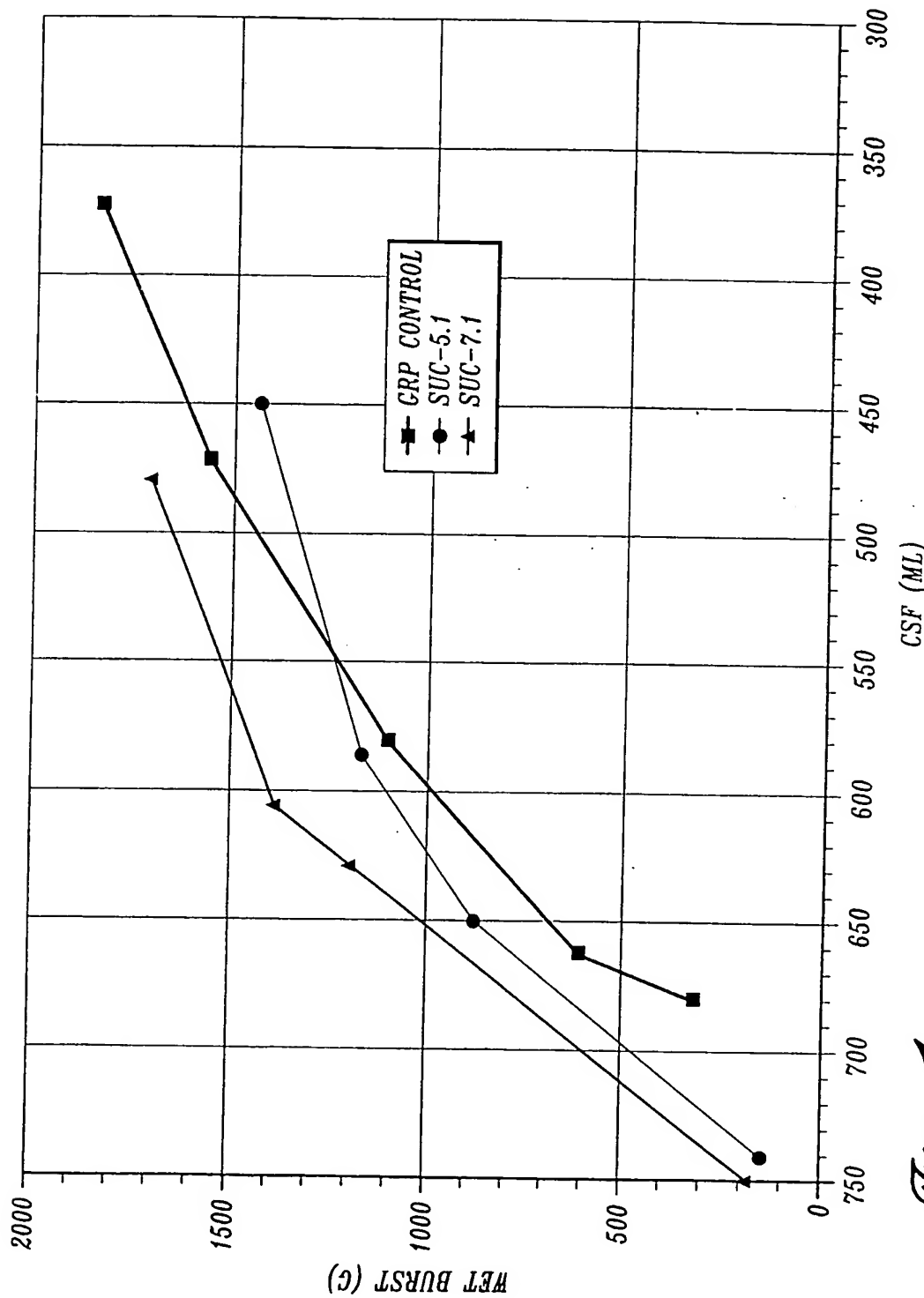


Fig. 1.

2/5

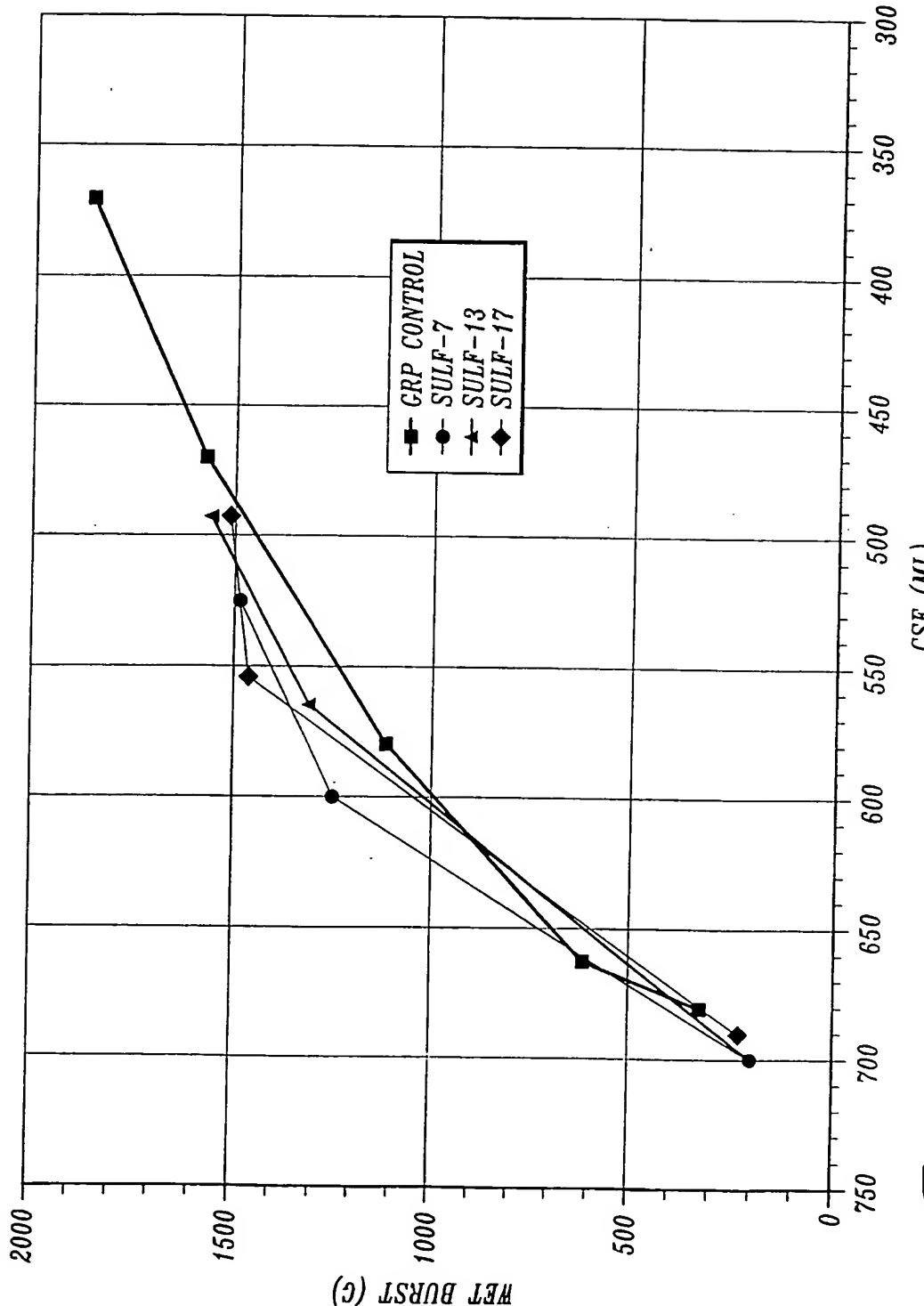


Fig. 2.

3/5

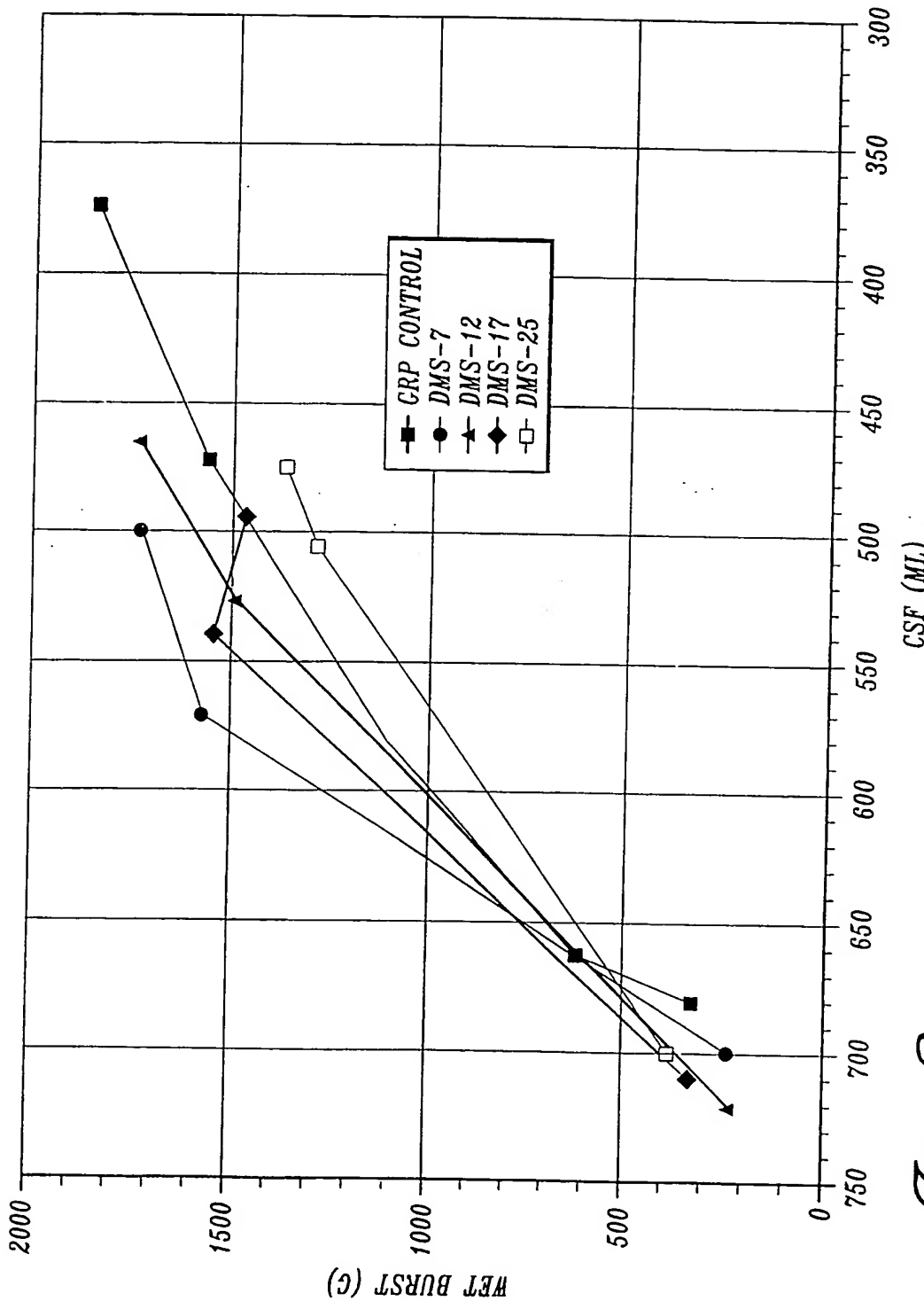


Fig. 3.

4/5

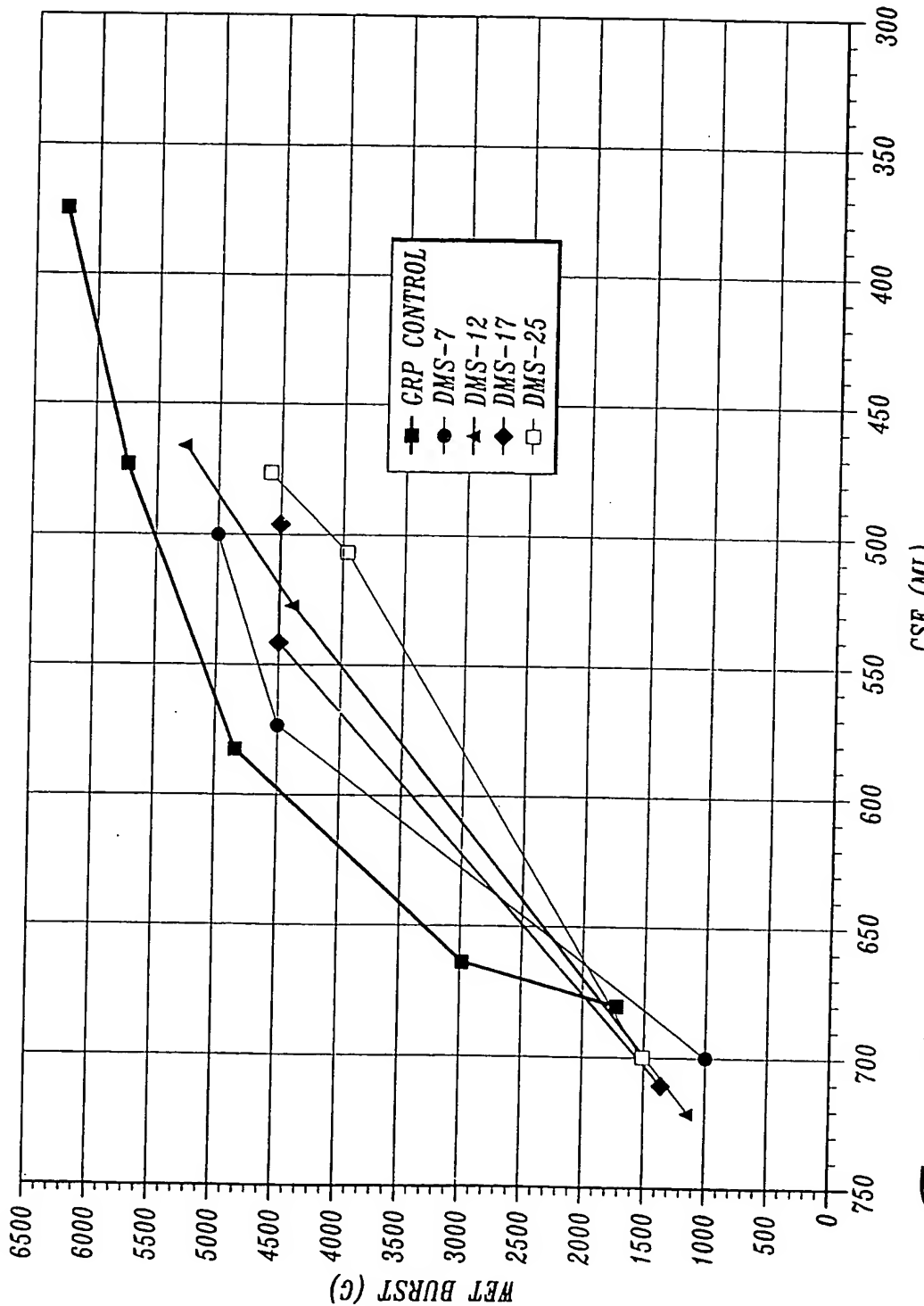


Fig. 4.

5/5

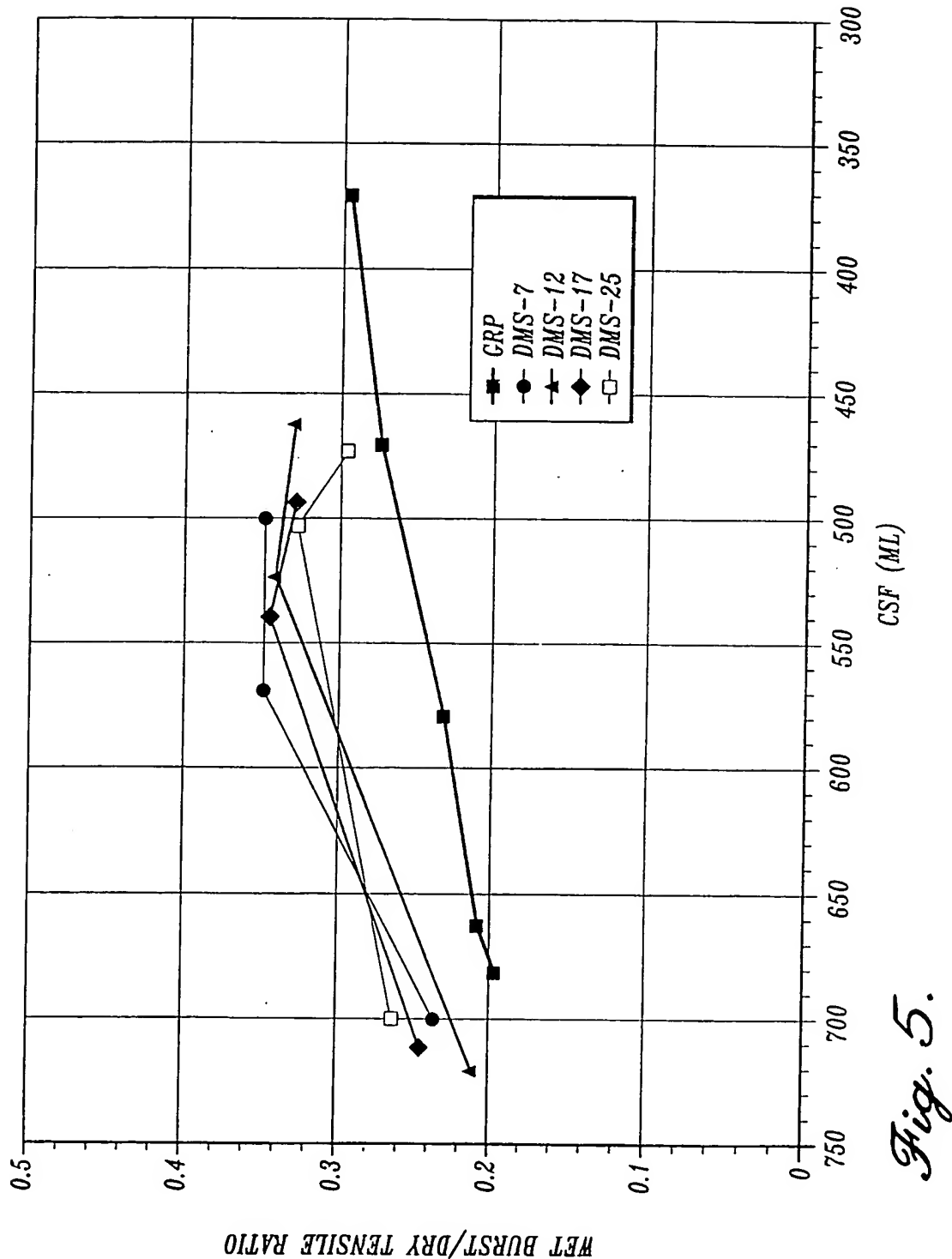


Fig. 5.

INTERNATIONAL SEARCH REPORT

Int. Appl. No
PCT/US 99/29884

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 D21C9/00 D21H11/20 D06M13/192 D06M15/263		
According to International Patent Classification (IPC) or to both national classification and IPC		
B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC 7 D21C D21H D06M		
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched		
Electronic data base consulted during the international search (name of data base and, where practical, search terms used)		
C. DOCUMENTS CONSIDERED TO BE RELEVANT		
Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 5 755 828 A (WESTLAND JOHN A) 26 May 1998 (1998-05-26) column 1, line 1 -column 7, line 50 --- Y	1,2,7, 9-13,22, 23,27, 29-32, 35-38, 41-43, 45,46 15,29 29
Y	US 5 447 977 A (HANSEN MICHAEL R ET AL) 5 September 1995 (1995-09-05) cited in the application column 39 -column 44; figure 1 --- -/-	
<input checked="" type="checkbox"/> Further documents are listed in the continuation of box C.		<input checked="" type="checkbox"/> Patent family members are listed in annex.
<p>* Special categories of cited documents :</p> <p>"A" document defining the general state of the art which is not considered to be of particular relevance</p> <p>"E" earlier document but published on or after the international filing date</p> <p>"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)</p> <p>"O" document referring to an oral disclosure, use, exhibition or other means</p> <p>"P" document published prior to the international filing date but later than the priority date claimed</p> <p>"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention</p> <p>"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone</p> <p>"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.</p> <p>"&" document member of the same patent family</p>		
Date of the actual completion of the international search	Date of mailing of the international search report	
10 May 2000	17/05/2000	
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl. Fax: (+31-70) 340-3016	Authorized officer Nestby, K	

INTERNATIONAL SEARCH REPORT

International Application No
PCT/US 99/29884

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	US 5 667 637 A (JEWELL RICHARD A ET AL) 16 September 1997 (1997-09-16) cited in the application the whole document	15
A		18-21, 25,26, 33,34, 39,40,44
A	US 5 137 537 A (HERRON CARLISLE M ET AL) 11 August 1992 (1992-08-11) cited in the application	
A	US 5 549 791 A (HERRON CARLISLE M ET AL) 27 August 1996 (1996-08-27) cited in the application	

INTERNATIONAL SEARCH REPORT

Information on patent family members

International Application No
PCT/US 99/29884

Patent document cited in search report		Publication date		Patent family member(s)	Publication date
US 5755828	A	26-05-1998	EP WO	0946810 A 9827262 A	06-10-1999 25-06-1998
US 5447977	A	05-09-1995	US	5308896 A	03-05-1994
			US	5614570 A	25-03-1997
			US	5807364 A	15-09-1998
			US	5998032 A	07-12-1999
			AU	5019893 A	15-03-1994
			AU	5019993 A	15-03-1994
			BR	9306920 A	12-01-1999
			BR	9306921 A	12-01-1999
			CA	2140263 A	03-03-1994
			CA	2140264 A	03-03-1994
			EP	0655970 A	07-06-1995
			EP	0655971 A	07-06-1995
			JP	8500270 T	16-01-1996
			JP	8500271 T	16-01-1996
			NZ	255814 A	25-06-1996
			NZ	255815 A	26-03-1996
			WO	9404351 A	03-03-1994
			WO	9404352 A	03-03-1994
			US	5693411 A	02-12-1997
			US	5672418 A	30-09-1997
			US	5547745 A	20-08-1996
			US	5641561 A	24-06-1997
			US	5607759 A	04-03-1997
			US	5547541 A	20-08-1996
			US	5611885 A	18-03-1997
			US	5789326 A	04-08-1998
US 5667637	A	16-09-1997	CA	2236415 A	09-05-1997
			EP	0858528 A	19-08-1998
			JP	11515063 T	21-12-1999
			WO	9716595 A	09-05-1997
US 5137537	A	11-08-1992	AT	134511 T	15-03-1996
			AU	649233 B	19-05-1994
			AU	6582090 A	16-05-1991
			BR	9005635 A	17-09-1991
			CA	2029014 C	18-03-1997
			CN	1053011 A	17-07-1991
			DE	69025548 D	04-04-1996
			DE	69025548 T	26-09-1996
			DK	429112 T	08-07-1996
			EP	0429112 A	29-05-1991
			ES	2083995 T	01-05-1996
			GR	3019165 T	31-05-1996
			JP	2834312 B	09-12-1998
			JP	3206176 A	09-09-1991
US 5549791	A	27-08-1996	AT	168421 T	15-08-1998
			AU	2606595 A	05-01-1996
			CA	2191567 A	21-12-1995
			DE	69503508 D	20-08-1998
			DE	69503508 T	28-01-1999
			EP	0765416 A	02-04-1997
			ES	2118604 T	16-09-1998
			JP	10503811 T	07-04-1998

INTERNATIONAL SEARCH REPORT

Information on patent family members

Inte. ornl Application No
PCT/US 99/29884

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
US 5549791 A	WO 9534710 A		21-12-1995

2/9/1

16095121

Basic Patent (No,Kind,Date): WO 200039389 A1 20000706

Family search of
WO 00/39389
2/14/01

PATENT FAMILY:
AUSTRALIA (AU)

Patent (No,Kind,Date): AU 200024804 A5 20000731

CARBOXYLATED CELLULOSIC FIBERS (English)

Patent Assignee: WEYERHAEUSER CO

Author (Inventor): JEWELL RICHARD A

Priority (No,Kind,Date): US 222372 A 19981229; WO 99US29884 W 19991216

Applic (No,Kind,Date): AU 2424804 A 19991216

IPC: * D21C-009/00; D21H-011/20; D06M-013/192; D06M-015/263

CA Abstract No: * 133(07)090921T

Language of Document: English

WORLD INTELLECTUAL PROPERTY ORGANIZATION, PCT (WO)

Patent (No,Kind,Date): WO 200039389 A1 20000706

CARBOXYLATED CELLULOSIC FIBERS (English)

Patent Assignee: WEYERHAEUSER CO (US)

Author (Inventor): JEWELL RICHARD A

Priority (No,Kind,Date): US 222372 A 19981229

Applic (No,Kind,Date): WO 99US29884 A 19991216

Designated States: (National) AE; AL; AM; AT; AU; AZ; BA; BB; BG; BR; BY; CA; CH; CN; CU; CZ; DE; DK; EE; ES; FI; GB; GD; GE; GH; GM; HR; HU; ID; IL; IN; IS; JP; KE; KG; KP; KR; KZ; LC; LK; LR; LS; LT; LU; LV; MD; MG; MK; MN; MW; MX; NO; NZ; PL; PT; RO; RU; SD; SE; SG; SI; SK; SL; TJ; TM; TR; TT; UA; UG; UZ; VN; YU; ZA; ZW (Regional) GH; GM; KE; LS; MW; SD; SL; SZ; TZ; UG; ZW; AM; AZ; BY; KG; KZ; MD; RU; TJ; TM; AT; BE; CH; CY; DE; DK; ES; FI; FR; GB; GR; IE; IT; LU; MC; NL; PT; SE; BF; BJ; CF; CG; CI; CM; GA; GN; GW; ML; MR; NE; SN; TD; TG

Filing Details: WO 130000 With international search report; Before expiration of time limit for amending the claims and to be republished in the event of the receipt of the amendments

IPC: * D21C-009/00; D21H-011/20; D06M-013/192; D06M-015/263

CA Abstract No: ; 133(07)090921T

Derwent WPI Acc No: ; C 2000-490857

Language of Document: English

WORLD INTELLECTUAL PROPERTY ORGANIZATION, PCT (WO)

Legal Status (No,Type,Date,Code,Text):

WO 200039389 P 20001012 WO DFPE REQUEST FOR PRELIMINARY EXAMINATION FILED PRIOR TO EXPIRATION OF 19TH MONTH FROM PRIORITY DATE

Inpadoc/Fam.& Legal Stat (Dialog® File 345): (c) 2001 EPO. All rights reserved.